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11 Publication number:

0 038 885

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EUROPEAN PATENT APPLICATION

21) Application number: 80301363.0

22 Date of filing: 25.04.80

(6) Int. Cl.3: C 08 J 9/28

C: 08 J 5/18, G 03 C-1/76 G 11 B 5/70, B 01 D 13/04 C 08 J 5/04, H 01 L 21/31

Date of publication of application: 04.11.81 Bulletin 81/44

(84) Designated Contracting States: BE DE FR GB NL (1) Applicant: Exxon Research and Engineering Company P.O.Box 390 200 Park Avenue Florham Park New Jersey 07932(US)

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64 Microcellular heterocyclic polymer structures.

(5) Highly useful novel microcellular polymeric structures, especially films and fibers, are prepared from certain solid polymers. Aromatic polysulfones, polyimides, polyhydantoins, polyamides and polyparabanic acid are the preferred polymers for the novel structures of the invention.

P 0 038 885 A1

1 The technique of casting polymeric articles is old and well established. In general, there are two 2 approaches to casting so One of these is casting either 3 a monomeric or partially polymerized syrup into a mold 4 or shape and conveying this into an oven or autoclave in 300 \$\omega\$ 5 gorder to finish polymerizing the article with a tempera- are 3 ture treatment. The other general type of casting involves solution casting which is also a long utilized 8 technique for producing plastic film and sheet materials. 9 The general technique of solution casting in-10 volves forming a solution of the film-forming polymer a suitable solvent, casting the resulting solution on a 12 suitable substrate, evaporating the solvent and winding 13 the resultant film on rolls. 14 Usually solvent recovery systems are employed 15 in order to recover the solvent and minimize the loss of 16 an expensive process component. 17 Solution cast opaque films have been conven-18 tionally prepared by adding pigments, fillers, flame re-19 tardants and solubilizers to a solution of the film-forming 20 material, which pigment acts as an opacifying agent. 21 out such an agent, such film would be colorless or trans-22 parent. Opacifying agents often embrittle the film. 23 Various processes have been described in the art 24 for preparing opaque films which rely for opacity upon 25 the presence of a large number of voids in the film. 26 Such films may be prepared by depositing a film from an 27 emulsion, i.e., either an oil-in-water or a water-in-oil 28 emulsion. 29 When a water-in-oil emulsion is used--i.e., 30 one in which minute droplets of water are dispersed in 31 a continuous phase of a film-forming material -- the emul-32 sion is deposited as a coating and the organic solvent 33 which comprises the continuous phase of the emulsion is 34 evaporated therefrom. This causes the gelation of the 35

- 2 -

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film-forming material and entrapment of the dispersed
1
   water droplets. The water is then evaporated leaving
2
   microscopic voids throughout the film structure.
             Still another technique for obtaining a porous,
4
    opaque, nonpigmented film is set forth in U.S. Patent
5
   No. 3,031,328. Basically, this process contemplates
6
    preparing a solution of a thermoplastic polymer material
7
   in a mixture of a volatile organic solvent and a volatile
9 - non-solvent liquid which has an evaporation rate sub-
    stantially less than that of the solvent. The clear
10
    homogeneous solution is then coated on a suitable backing
12 material and dried by evaporation to produce an opaque
    blushed film which is adapted to be rendered locally trans-
    parent by heat or pressure. These films are useful in
14
    recording films.
15
              Other techniques for forming opaque, porous,
    nonpigmented, microporous thermosetting films are set
    forth in U.S. Patent No. 3,655,591.
18
         Nevertheless, in spite of the above, the art has
19
    never appreciated the unique articles which result when a
 20
     specific type of polymer is cast in a certain manner to
 21
     produce an essentially nonporous, nonfoam microcellular
 23 structure which has unique and unusual properties and is
                                                              22
     incidentally opaque. The art has concentrated on tech-
                                                              . .
     niques wherein the opaqueness is the sine qua non of the
 25
     structure and the other properties are not if signifi-
 26
 27 - cance: A y Properties of Kor militarrole
         In accordance with the present invention, unique
 28
     microcellular, nonporous, nonfoam polymeric articles such
 29
     as films and fibers are prepared by a novel solvent/non-
 30
     solvent casting technique.
 31
 32 ... It is known that films, fibers and other struc-
 33 tures can be made out of solvent cast polymers such as
 34 those described in U.S. Patent No. 3,661,859.
     ticular polymers are referred to a 1,3-imidazolidene-2,4,5-
 36 trione-1,3-diyl. The repeating heterocyclic ring structure
     of these polymers is shown as follows:
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    in a may 000, to relative or anic sulvant und a volatile
7
    These and similar polymers and processes for their prepara-
8
    tion are disclosed in U.S. Patents 3,591,562; 3,547,897;
9
    3,661,859; 3,684,773; 3,637,843; and 3,635,905
10
       Other preferred polymers are aromatic polyamides, 1
11
    aromatic polysulfones, and polyhydantoins which have been
12
    described in the art. See, for instance, Netherlands
13
    6809916, Belgium 723,772; German 1,807,742; 1,805,955;
14
     1,812,002; 1,812,003; and 1,905,367. Polyimides are well-
15
    known and are described in such publications as British
 16
     1,240,665; U.S. 3,486,934; U.S. 3,536,666; French
                                                              3:
     1,488,924; French 1,549,101; Russian 218,424; German
 18
     1,301,114; Netherlands 7,001,648 and the like.
 19
        The detailed preparation of these polymers and
 20
     solutions of these polymers in suitable solvents are set
 21
     forth in the above-recited patents and others also in the
 22
     art, and therefore need not be repeated here except as is
 23
     necessary to understand the invention.
 24
               The preferred microcellular structure from the
 25
     polymers of the invention are characterized by high temp-
 26
     erature thermal stability, organic solvent resistance,
 27
     relatively high tensile modulus, tensile strength and
     ultimate elongation with low shrinkage at high tempera-
  29
     tures and are slow smoke formers when ignited.
  30
        Nonmicrocellular film from the preferred polymers
  31
     have relatively high dielectric strengths. These proper-
      ties have been found to offer outstanding advantages when
      used as films in flexible circuitry, for use in auto 25 1.
  34
      air bag circuits, light monitoring circuits, and telephone
```

circuits because of their ability to be soldered. They also are useable for fibers, where high tenacity and modulus 2 are required. 3 ភេស ស្ពេងសៀមមាន ខែជា ខ្មែ**មកំណា**លប្ផុំ**ងគ្ ខ្**កួត្តិសំពេលប្រុស្ស However, in these applications, the structure 4 . has a relatively high cost per unit of weight. 5 It would be desirable to have a structural article possessing es-6 sentially the outstanding properties of the above-described 7 . noncellular film so that it can be used for the applica-8 tions listed above, but it would be less dense. 9 ducts of low density and still superior properties could 10 be obtained, it would mean that a novel new structure of 11 outstanding cost-performance utility would exist. 12 It has been discovered and forms the fundamental 13 substance of the invention that such relatively low density 14 microcellular structures can be prepared and are novel 15 themselves. These films are very thin and are essentially 16 nonporous, i.e., they are made up of closed cells. The 17 18 technique of preparing them forms a portion of this in-19 20 If the advantages delineated above for the lower density material were all that the material contributed, 21 its existence would be welcomed and its utility would be 22 considered outstanding. Notwithstanding the outstanding 23 utility of the lower density material, it has been dis-24 covered that the material has additional unique properties 25 of its own which make it extremely valuable in addition 26 to those properties enumerated above. 27 The preferred polymer species of the invention 28 are polymers of 1,3-imidazolidene-2,4,5-trione, i.e., 29 polyparabanic acid, herein referred to as PPA; and poly-30 (imino-1,3-phenyleneiminocarbonyl-1,3-phenylenecarboxyl), 31 hereinafter referred to as IPP. The particular conditions, 32 reagents and uses are especially well-suited for the PPA 33 or IPP polymers and structures resulting therefrom. 34 theless, it must be emphasized that other polymers of this 35

invention can be handled in an analogous manner to make

```
structures which have some similar properties. These
2 latter include the soluble polyamide-imide, polyimides,
                polysulfones, polyamides, and various soluble polyhydan-
                toins.
                                                         In general, the polymers of the invention will
               be comprised of sufficient repeating units to be solids
                at room temperature. The repeating unit can contain
                heterocyclic rings.
 8
  9
                                                         The heterocyclic ring will be 5-membered and
                will contain carbon, and nitrogen linkages wherein at
                 least two of the carbons will be in the form of carbonyl
                groups, i.e., O which are separated by a nitrogen atom.
                                                           THE COURSE HARD MALE AND AND A SECOND
                         Examples of heterocyclic rings which fall in
                 this case are: Park the maker of the tender of the control of the 
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                                                                                                                               Poly(hydantoins);
                                                                              rac{1}{2\pi}R . In the field r , we have r
     25 , (oxed{2\pi}_0 \omega_0 \omega_0 ) ໄດ້ rac{1}{8} າ ຄືຊື່ 48 , ພາຍໄຈ ການພັກ ເປັນປະຊານ ພ້າການ ໄດ້ການ ເຊື່ອ ຄົວ ເພື່ອ oxed{2\pi}_0
     ner ingried og færreð tæ es 125. Edmi þer litte ner til lænni.
                  భినాక్ ఉన్నా గురు, కంగు మండు, అమేమ్కు గుమ్మెమ్డిశానగా ఈ చాయానా నాని మీకి ముందుకు ముందుకు కారా
                                   లాక్ష్మార్డ్ మాట్లు <mark>రాజాలు క్రామాన్నారి. క</mark>్రామ్ ఇవ్వార్లు కార్యాల్లో ఉన్నారి. క్రామాన్ని క్రామాన్ని క్రామాన్ని
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           24.00 Other preferred polymers have repeating units
               as follows: 310 dur tide to this part of the tide is
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                                                                                                                                               -Polysulfone
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                    Markanings . While A . X was Or / OR = Aromatic or
                                     in the victoria and
 18
                                                   Wherein Z is a number from 20 to 1,000, prefer-
                ably 50 to 200.
  20
             Although casting in general is a relatively
                well-known process, for each polymer and solvent system
  22
                there are unique problems brought about by the particular
  23
                 solvents which must be used and the properties of the
  24
                polymer itself. Very generally, PPA's are soluble in
  25
                 moderate hydrogen bonding dipolar, aprotic solvents.
                 This presents a practical problem in casting, since
  27 .
                 solvents which are available at a reasonable cost have
   28
                 relatively high boiling points and are of low volatility,
   29
                 except at relatively high temperatures. The effect of
   30
                 these parameters is that when PPA is cast into even
   31
                  relatively thin structures, a film, for instance, it is
   32
                  relatively difficult to remove the last small amounts of
    33
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3 4 3 3 7 6 3

solvent from the structure, e.g., film. 1 For instance, dimethylformamide (DMF) is con-2 sidered to be one of the best solvents for working with 3 PPA solution formulations. Tt boils at 156°C and its excellent solvating effect results in the fast dissolu-5 tion of PPA along with the formation of low viscosity 6 solutions. 7 Nevertheless, this combination of low volatility 8 and high solvation, which characterize a good solvent makes 9 the removal of the last amounts of solvent from even thin structures such as films very difficult. Therefore, film 11 casting processes must be conducted with extremely high 12 drying temperatures in order to get good solvent removal 13 at reasonable production rates. 14 In accordance with this invention, low density 15 nonporous, microcellular film structures, e.g., PPA, 16 aromatic polyamides and others listed above are prepared 17 by first solvent casting of film. Prior to complete drying, one precipitates the film by contacting the film with an antisolvent, such as water. A basic requirement 20 for the antisolvent is that it be miscible with the solvent 21 in the polymer solution. 22 More specifically, the inventive process involves 23 the steps of: 24 preparing a casting solution of the polymer; (a) 25 casting a wet film onto a surface or ex-26 truding a fiber; 27 (c) paritally drying the cast film or fiber; 28 (d) contacting the wet film with an antisolvent 29 The traction of the second second such as water; and 30 (e) removing solvent and antisolvent by com-31 pletely drying the now nonporous, microcellular 32 article. ా ఎక్కార్లు కృత్తున్న అక్షాణాడు. ఇద్ది కారాజికేస్తా 33 In order to obtain a film with very low density, 34 one can eliminate steps (c) and for step (d), expose the film to an atmosphere of high humidity.

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Additives, such as flame retardants, oxidation
1
    inhibitors, plasticizers, etc., should be dissolved in
2
   the solvent with the resin prior to casting. So cities
3
            The solvents which can be employed in accordance
4
    with this invention are moderate hydrogen bonding dipolar,
5
    aprotic solvents. These solvents have been described in
6
    U.S. Patent No. 37661,859. The preferred solvents are
7
    N, N-dimethylformamide, N-methylpyrrolidone, N, N-dimethyl-
8
    acetamide and dimethyl sulfoxide.
9
           .g. The antisolvents as mentioned above must be
10
    miscible with the solvent. Typical of the antisolvents
11
    are water, aliphatic alcohols such as methanol, ethanol,
12
    propanol, butanol, and the like; aliphatic ethers such
13
    as methyl ether, ethyl ether, methyl ethyl ether, propyl
14
    ether, methyl propyl ether, ethyl propyl ether and the
15
    like; and aliphatic ketones such as acetone, di-ethyl
16
    ketone, methyleethyl ketone and the like. The preferred
17
    antisolvent is water.
18
           The concentration of the resin in the solution
19
20 should be such as to not produce a viscosity which would
21 make the solution too difficult to handle. Typically,
    the suitable viscosities can be determined by simple ex-
22
    perimentation of the fact that the above of their
23
       Generally, for ease of operation, the concen-
24
    tration of the resin in the casting solution may be such
25
    that the Brookfield viscosity at 25°C is between about 80
26
    and about 800 poises. Desirably, the viscosity for the
27
    greatest ease of operation can be between about 200 and
28
    300 poises. - The contract of the second
29
       Prior to casting, it is desirable to filter the
30
    casting solution so as to remove any trash and gel par-
31
     ticles. The transfer of the same
                                    ాగు జూన్తి చారుకోందానికి ముందుకోంద
32
        In general, there are two methods according to
 33
     the invention that can be used at step (d) described above
 34
     to form the novel cellular articles of the invention.
 35
     These are: 3 - mail or many earlier and a con-
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محول المسلومات المسلوبينية المالية المالية

(a) Method 1 - The wet film or fiber, is 1 exposed to an atmosphere of high water humidity, 2 followed by a direct water washing, followed by its off 3 drying. As is true of all of the techniques, 4 the thickness and shape of the structure is con-5 introlled by its original casts or extruded thick-listing 6 ness and shape and solids content. Precipitating . 2.5 7 the structure in a high humidity environment and the M 8 rather than initial direct water contact is im-the contact 9 portant. The reason is that too rapid precipi-10 tation and solvent removal will cause wrinkling 11 of the structure, which is very undesirable. 12 13 film or fiber, is obtained by solvent casting 14 or extrusion; then it is partially dried to a 15 greater or lesser extent. This serves two 16 purposes; prevents wrinkling and increases the 17 density of the structure. Then it is water 18 washed and dried completely. The density will 19 vary according to the amount of solvent removed 20 in the initial drying step. The state of the state of the 21 rest for the rest density of the second could be possibly relatively to the Method 1 gives a density of about 0.45-g. per-22

cubic centimeter, Method 2 gives a density varying from about 0.3 to 1.5 and preferably 0.3 to 1.2 g. per cubic centimeter. When operating in the solution casting mode, the following considerations will be pertinent.

23

24

25

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Density is largely dependent on the weight fraction of polymer in the wet film at the instant precipitation occurs. Casting solutions of PPA ranging above 30 weight percent resin are difficultly handled in conventional solvent casting equipment due to their very high viscosity. The high viscosity solution can nevertheless be readily obtained through extrusion through an appropriately shaped die.

The method 1 technique contemplates the use of 36 the most viscous solution that can be handled, i.e.,

37

20 to 50 weight percent PPA depending upon molecular 1 weight of the polymer. 2 Method 2 permits the use of a more dilute solu-4 of tion with its concomitant easiers handling advantages, and 5 m relies upon the evaporation of more solvent from the film after it is cast and prior to first precipitation. 6 ा हुल हुन् हुन् This callows a wider range of densities that can ् 7 be obtained by Method lims mandages such at man and section for a 8 The practical limit which sets the maximum den-9 sity which can be obtained in Method 2 is the minimum ! : : 10 amount of solvent which must remain in the polymer in order 11 for precipitation to occur when the cast polymer solvent. 12 structure is contacted with water or other antisolvent. 13 The minimum density of Method: 2% is limited by 32 14 the maximum amount of solvent that can be left in the wet 15 film at the precipitation step, which will not cause sur-16 face wrinkles on the film or fiber surface. This will vary 17 depending on the choice of anti-solvent as can be simply 18 determined. He resummed a codano libor establica a maligne above o 19 The range of densities can be further increased 20a 21: by: (a) calendering the resulting cellular film, (b) to the orienting the film and the fibers to elongate and reduce 22 the volume of the cellular portions, or (c) the use of 23 different mechanical equipment designed to handle the ex-24 tremely viscous polymer solutions, for example slot ex-25 truders. That latter approach would increase the density 26 of the microcellular material approximately proportionately 27 to the amount of solvent reduction in the original polymer 28 solvent solution. Thus when solution extrusion equipment 29 is used, much higher polymer solvent contents can be 30 handled as compared to the casting methods described above. 31 In accordance with this invention, the nonporous 32 microcellular structures can be prepared as a free film, 33 a permanent coating on a surface or as an impregnating 34 substance. grant went to the grant of infer and a grant of the 35 The free film is prepared by laying down a layer. 36

of casting solution onto the desired flat surface which

ration from the control of the same properties of the field of the fie

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conveys the wet layer sequentially into a drying zone
1.
       sonas to partially dry the film, a water zone (bath, we have
2
       vapor or#spray) rands to aufinal drying zone. @Suitable sur-
3
        faces which may be employed are metal, which has been the
4
       polished or embossed, chrome plated metals, release paper
5
        and others known in the casting art. When needed, a re-
6
        lease agents can be included in the casting solution to
7
        facilitate removing the finished film from the casting
8
      surface. The indicated above, so as to obtain a very low -
9
        density film; the first drying step can be eliminated. The
10
              ್ಯಾಕ್ಸ್ ಫ್ರೌಫ್ g Suitabled equipment: for laying down the wet ಕರ್ಮ ಅ
11
        film of casting solutions are casting boxes, reverse roll
12
        coaters and pressured extrusion dies. The choice dependes
13
        upon the thickness of wet film to be laid down and the
14
15 viscosity of the casting solution. With casting solu-
        tions having a Broodfield viscosity less than 200 poises
16
        one can employ a reverse roll coater. Intermediate vis-
17
        cosities and wet film thickness are handled best by cast-
18
        ing boxes. The typical ranges are Brookfield viscosities
 19
        of 100 to 300 poises at wet film thickness of 10 to 30 10
 20
        mils. Very high viscosities, 300 to 1000 poisés require
 21
        extrusion dietequipment. The the transport of the transpo
 22
          The wet film exposed to the first drying zone
 23
        will have the initial composition of the casting solution.
 25 CAfter partial drying or in the absence of the partial
         drying step, the solvent content of the film should
 26
         preferably be between about 20% to 50% by weight. Never-
 27
         theless, greater or lesser percentages may be employed.
 28
         However, if less than 20% solvent is present, the desired
         cellular structure will be formed in the water zone slowly
 30.
         and the cellular structure will not be as perfectly formed.
 31
         If the residual solvent is greater than 50%, the rate of
 32
         water absorption will be too rapid causing the film to be
 33
         deformed. In between a well-formed cellular structure will
  34
         be produced with little or no film deformation.
  35
                            The water zone may consist of water vapor, water
  36
         spray, a water bath, or any combination of these as long
  37
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as water is rapidly absorbed by the film. In addition to
 1
     water being absorbed into the film, solvent is extracted
 2
   from the film such that the solvent content of the micro-
     cellular film should be less than 10%. During the final
     drying step, the microcellular film will tend to melt with
    loss of microcellular structure if the solvent content is
     much greater than 10%.
   As the film is partially dried and as it is con-
    tacted with water, the film thickness and cell diameter,
     will decrease due to removal of solvent.
              The film is dried in a final drying zone which
     preferably is a stage zoned oven having temperature gradi-
     ent of from 175 to 270°C. The final solvent content is
 13
     usually < 3000 ppm. A further reduction in film thickness
     and cell diameter occurs during the final drying.
 15
          The properties of the finished microcellular
 16<sup>74</sup>
     film are basically determined by (1) film thickness,
17
     (2) film density, and (3) cell diameter. It is apparent
18
    that final film thickness is substantially less than that
19
    of the initially cast wet film since as indicated above
    the thickness is reduced in each step of the casting
    operation. The largest reduction must occur in the first
    drying zone since from 40% to 75% of the solvent is de-
23
    sirably removed. Thus, if a particular thickness of fin-
24
    ished microcellular film is to be obtained, the initially
25
    cast wet film must be from 2 to 3 times this thickness.
26
    The upper practical limit of thickness of the finished
27
    microcellular film of this invention is about 20 mils.
28
    This is to be contrasted with foams which are formed by an
29
    expansion process. Thus, the thickness of the foamed ar-
30
    ticle and the cell diameter increase during the foaming
31
    process. This imposes a practical lower limit of greater
32
   than 20 mils thickness for foamed articles.
33
                                                The lower
   practical thickness for these microcellular film is about
34
   1 mil which is far below that attainable by foaming.
35
   Hence, as a practical matter, the films of this invention
36
   range from about 1 mil to about 20 mils in thickness.
37
```

图图2600

1 Film density is governed by the volume fraction of the microcells. For example, if the volume fraction is 3 0.50, then the density of the microcellular film will be 50% of that of dense film from the same resin. The density of microcellular film has a practical lower limit of about 30% of the dense film value. The upper limit is about 90% of the dense film value. Foams by contrast are usually less than 30% of the corresponding dense article. The range of cell diameters usefully employed in accordance with this invention can be from about 0.1 to about 10 microns. Although cell sizes can be greater or lesser, for most useful applications, the smallest cell diameter is preferred such as from about 0.1 to about 5 microns, although there are some exceptions to this rule. In any event, the mechanical strength, compressibility and toughness are increased the smaller the microcells are for any given film density. By contrast, foamed articles 17 usually contain cells having greater than 10 microns diameter and their mechanical strength and toughness are low. 19 The shape and distribution of the microcells are also im-20 portant. In general, microcellular articles, however they may be made, have structures made up of either open or closed cells. All of the film of this invention contain predominantly isolated spherical closed cells which have 24 such a uniform distribution that only occasionally do two cells impinge on one another. By virtue of the essentially discrete spherical closed cells, the transmission 27 of gases, vapors and liquids are so slow by the film that it can be considered to be impermeable when compared to semi-permeable microporous membranes and foams. 30 Cellular film made from PPA solutions in DMF 31 were prepared on equipment which is normally used to make porous cellulose acetate film for electrophoresis appli-33 cations. The equipment consisted of a casting box applicator, a sixty-foot continuous stainless steel belt, and 36 four chambers equipped to control humidity, temperature and the rate of air flow.

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Provisions were also incorporated to spray water
 1
     onto the moving continuous belt for the purposes of initial
 2
     precipitation and for washing the solvent from the film.
 3
     The first technique used was that described for
 4
     Method 1. The polymer was PPA in 20% concentration in DMF.
 5
     The humidity was controlled at 90-95° in the first zone.
 6
     The film initially precipitated due to absorption of water
 7
     vapor. Additional precipitation and solvent removal was
 8
     effected by direct immersion in a water bath following by
 9
     drying! & this or about the entire from a depocatific
 10
     Under these conditions of high humidity, the wet
 11
     film absorbs water vapor rapidly due to the hydroscopic
12
    nature of the DMF solvent, but the absorption is much slower
    than if the film were directly immersed in water. Films
14
    having a uniform cellular structure were obtained having
    densities of about 0.45 g. per cubic centimeter.
     The belt speed varied from about .50 feet per
17
    minute to about 2.5 feet per minute. The temperature was
18
    about 100°F. The air rate was about 900 to 1,300 cu. ft.
19
    per minute. The thickness of the wet film varied from
20
    about 8 mils to about 20 mils. The total time in the oven
21 -
    ranged from about 6 minutes to about 15 minutes. Generally
22
    time periods above 10 minutes and less than 20 minutes
23
    appeared to be satisfactory.
24
          Subsequent work has been done to produce cellu-
25
    lar films which have relatively high densities, i.e., up
26
    to 1.1 gm/cc.
27
              For this, the above-described equipment was
28
    modified somewhat to permit the use of a Method 2 type
29
    approach. It required the addition of heaters to the
30
    first chamber in order to provide for some gradual ini-
31
    tial solvent removal, which is a step required to control
32
   the increase in the density of the film and to prevent
33
   wrinkling. Equipment to spray water onto the moving stain-
34
   less steel belt was installed in the second chamber to
35
   precipitate the film. Water sprays in the third and fourth
36
   chambers were provided in order to wash out additional
37
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solvent, e.g., N.N-dimethylformamide (DMF) prior to stripping the film off the belt and subsequent drying. Passage of the solvent loaded film into an ir-3 regular water interface produced a nonuniform surface on the precipitated film. Therefore, an air knife was in-5 stalled which directed an air flow downward onto the sur-6 face of the belt and provided a relatively uniform water. 7 interface for the wet film to pass into the balls 8 Although useful microcellular articles in par-9 10 ticular foams and semipermeable membranes have thickness >0.020" have previously been known, such structures have 11 not been found to have utility as a high performance en-12 gineering material in thinner gauges. By contrast, they 13 tend to be of relatively low strength. For the first time, 14 the microcellular film of this invention provide materials 15 in thin gauges which combine the properties necessary for 16 high performance engineering materials with some of the 17 desirable characteristics of light weight microcellular 18 materials. Some of the more important properties are il-20 lustrated in Table I where microcellular PPA-M is compared 21 to several commercial foams and dense film made from engineering plastics. The aromatic polyamides, polyimides, polyhydantoins, polysulfones and polyamide-imides of this invention have properties similar to the PPA films. to the process of the constant of the constant specific such

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,			Density (gm/cc)	Tensile Strength	Elongation (%)	Tearing Strength (gm/m11)	Dielectric Constant (103 cps)	St	(KV/m11)	. Q		(a) Properties from "Modern	(b) Property not 11sted.		- :			
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The above data show that microcellular PPA-M film is com-1 petitive with a widely used noncellular film for en-2 gineering thermoplastics, in tensile strength, percent 3 elongation, tearing strengths and dielectric strengths. On the other hand, the dielectric constant of the micro-5 cellular film of this invention is comparable to the very 6 structurally weak foamed sheetings. The very low di-7 electric constant is of great utility when used in elec-8 tric insulation for both power and signal transmission. 9 Thus, the microcellular film of this invention provides 10 a dielectric insulation which has a desirably low di-11 electric constant and at the same time, high dielectric 12 and mechanical strengths. Semipermeable membranes, 13 although not used for structural or insulation appli-14 cation, would be even less useful than foam in these applications due to their high porosity and low strengths. 16 Although it is predictable that mechanical 17 properties such as modulus and tensile strength will de-18 crease with decreasing density, it was found that these 19 mechanical properties were not sufficiently diminished 20 to seriously affect the utility of the cellular article for many applications. Moreover, in the case of the film, 22 the propogating tear strength was better than that of .23 the dense film. 24 The dielectric constant will decrease with 25 decreasing density, and therefore, the dielectric constant 26 for the cellular products is lower than that for the 27 dense film products. This makes the cellular film more 28 attractive for use as insulation, e.g., for microwave circuitry, especially where transmission is to be over 30 relatively long distances. In an analogous fashion, the 31 lower thermal conductivity makes these structures de-32 sirable for thermal insulation. 33 As is the case of the dense film, the cellular 34 film also withstands commercial solder bath temperatures, 35 i.e., 500°F vs. < 300°F for foamed sheetings.

```
1 or One important and highly advantageous property of the
  cellular film, as opposed to the dense film is that copper
  Ecircuits can be electroplated directly onto the cellular
  film, with much higher peel strengths for the electroplated
   copper, on the cellular film than on the dense film.
5
    For example, peel strengths for copper electro-
6
   deposited on the dense film were in the range of about
    2.5 to 3.0 pounds per inch. But peel strengths on copper
    electrodeposited onto the microcellular film were in the
   range of about 8 pounds per inch.
     This is an extremely important aspect of the
11
12 cellular film which gives it an outstanding advantage,
    taken in combination with its other properties, over dense
    film. But the first the care to a most roads to a market a well
15 Not only are the adhesion values exceedingly
16 high for the electroplated copper circuit on cellular
17 film, and also laminates prepared with adhesives but the
18 use of the cellular film permits the omission of a bother-
19 some process step. Thus, before copper laminated onto
20 plastic films which normally contain small quantities of
    absorbed water can be soldered, the unit must be dried to
21
    remove absorbed water. If it is not, the absorbed gater
    tends to be driven from the film during the soldering op-
23
    eration, because of heating of the composite unit. This
24
    rapid generation of steam causes the copper to be de-
    laminated from the film substrate.
26
              When the cellular, film of the invention is
 27
    utilized, the copper does not delaminate. It is theorized
    that this is due to the fact that there are numerous micro-
    cellular voids into which the water can expand rather than
 30
    escaping through the surfaces. Therefore, delamination
 31
     is effectively prevented. This is an exceedingly useful
 32
 33
     property.
              These films are much more flexible than, dense
 34
     film of the same thickness, which is an advantage for thick
 35
     multilayer structures.
 36
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Another important feature of the structures of 1 the invention involves selective surface etching by strong 2 3 bases or acids. This removes the film covering the microcells, either completely or in any pattern desired. The 5 exposed microcells, can then be electro or chemically 6 plated with far better adhesion. In fact, grooves can 7 be etched into the surface into which conductive metals can be deposited with excellent adhesion to the exposed 8 9 microcells and with excellent separation and insulation All in all, cellular film, because of its com-11 bination of properties and its relatively low cost, is 12 an ideal material for flexible circuits and flat conductor 14 scables. to community of module the made religions of above The preferred structures produced by the practice 15 of this invention are characterized by the presence therein 16 of a large number of discrete closed cells. Substantially 18 all of these cells or voids are less than 25 microns, and 19 preferably less than 2 microns, in size. Most preferably the cell size is less than 5 microns. The average cell 21 size and cell size distribution is governed by the condi-22 tions under which the structures are made, e.g., tempera-23 ture solvent, antisolvent, polymer solids content of cast-24 ing solutions, etc. The range obtainable is from about 25 0.1 to 25 microns. The column to Line was a large with the second of the column to Unless some color-forming material has been in-26 cluded in the composition, such as a soluble dye, the 27 preferred films of this invention are opaque and off white. 29 Colored films may be obtained by incorporated small amounts of dyes. A film having an apparent thickness of, for examples, 10 mils will have a real solid thickness which is equal to the sum of the thickness of each wall between the discrete cells lying along a path perpendicular to 34 the outermost planar surface of the film which may be, 36 for example, no more than 3 mils. Thus, the film is of

l sufficient apparent thickness to provide the required amount of strength.

Furthermore, the diffusion per unit of time of

4 a vapor or a liquid through a unit area of the films of

this invention is far smaller than semi-permeable mem-

. 6 g branes: A the lamin beforebilled general

5 . . .

7 - The compositions of this invention are particu-

8 larly useful when precipitated onto fabrics made from fiber

9 glass, resinous yarns, vegetable or cellulosic yarns and

10 cords. When these fibers or cords are coated with the

11 structures of this invention, an opaque or whate fabric

12 is obtained without the addition of pigments as needed

13 in the fabric heretofore employed. These coated fabrics

14 have very desirable flexibility. The fact that pigments

such as TiO2 are not needed to obtain whiteness in fibrous

fabrics is quite significant since this has been a problem

in the art due to the adverse effects these pigments have 17

18 on the resulting fabrics. For example, it is known that

pigments such as TiO2 weaken the tensile strength of the

20 fabric.

21 The fibers may be coated with the compositions

of this invention by either of the first two methods des-

23 cribed above. One method found to be suitable is to dip

24 the fibers into a solution which contains resin, solvent

and nonsolvent in amounts indicated hereinabove. Upon

precipitation a fabric having the desired whiteness and

softness is obtained without the addition of pigments such

as TiO2 28

Although the above discussion has been made with

30 reference to films as discrete articles, it is to be noted

31 that films in terms of surface coatings with unique and

important properties and which are bonded to a substrate

33 can also be produced according to the technique of the

34 invention.

The structures of this invention may be formed 36. as surface coating films by either of the techniques

described above. Thus, they may be applied by extrusion, 1 brushing, spraying, dipping, roller coating, or knife 2 coating followed by precipitation and drying.

The compositions of this invention are particularly useful when employed in spray applications. As already indicated, compositions of this inven-6 tion may be applied as films to various types of surfaces or substrates. These surfaces may be of the type whereby g the film is to be removed by a suitable method or of the 10 type where it is adhered to the final substrate such as the metal of an automobile. Among the more suitable sur-11 12 faces which may be coated with the cellular structures of this invention are steel, treated steel, galvanized steel, concrete, glass, fabrics, fiber glass, wood, plaster board, aluminum, treated aluminum, lead, copper and plastics. The most preferred surfaces are metals such as treated steel and treated aluminum.

Films formed from the compositions of this in-19 vention may be air dried, vacuum dried or bake dried at 20 elevated temperatures. Although considerable emphasis has been placed on cellular film formation and applications, it is an improtant feature of this invention that cellular fibers of high strength can be produced utilizing the technique A profession of opening at a prefession of the invention. Fibers made by the conventional wet spinning 26 techniques of the art are never left in cellular form, 27. but are remelted and oriented in order to eliminate the cellular structure which gives rise to fibers having low modulus. In this invention, the polymers used have such 30 a high modulus that the microcellular fibers can be used with only moderate-orientation. Ordinarily, orientation is used to improve fiber strength. 33 This gives rise to a microcellular fiber which 34 can accept dyes readily. Moreover, the fiber has the 35 capacity to absorb moisture. Thus, it will be comfortable

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l in contact with the human body. The capability of ab-
 2 sorbing moisture is often the difference between synthetic
 3 c fabrics which may feel clammy and natural fabrics such
 4 as cotton, the latter being much more comfortable because
 5 of their water absorptive capacity. Permanent press fab-
     rics can be made due to the high softening temperatures
 7 of these novel fibers.
 8 The microcellular films, fibers and other struc-
     tures can also be electrocoated with various metals such
 10 as copper, aluminum, and the like in order to form thin
 11 conductive coatings with a minimum of coating metal.
 12 Electrocoated structures can be used in a wide
 13 variety of decorative and utilitarian applications. These
    involve automotive trim, under-the-hood uses, and radia-
 Electrodeposition and chemical metalizing can
 16
    also be used to coat catalytic metals such as palladium,
 17
    platinum, nickel, and the like within the interstices
 18
    of the structure so that it can be used to form an ex-
19
    tremely high surface area, artifical surface for conduct-
    ing catalytic reactions at relatively high temperatures.
22.
              The cellular structures of the invention are also
    highly useful for specialty applications where highly tena-
23
    cious painted surfaces are required.
24
             The invention is further illustrated by the
25
26
    following examples.
27
    Example 1
             Utilizing the technique and apparatus described
28
    above, a series of runs was carried out in the apparatus.
29
   A casting solution was prepared by combining 19.7 weight
30
   percent PPA, 79 weight percent DMF and 1.3 weight percent
   of octabromobiphenyl (which is an excellent flame re-
32
33
   tardant for PPA film).
34
             All PPA utilized in these examples was prepared
   from the reaction of HCN with diphenyl methane diiso-
35
```

cyanate.

The state of the s

Three rolls of wet film were cast onto a moving 1 belt from the prepared solution, and partially dried in a 2 3 Circulating air oven at 180°F. Upon exiting from the oven, the film was rapidly precipitated by spraying with water, 5 followed by immersion in a water bath and completely dried in a circulating air oven. A fourth film was cast as a control but was not <u>;</u>7 subjected to the water bath so as to obtain conventional 8 9 noncellular product. has accommend the fact The resulting level of octabromobiphenyl in the 11 dense film was about 6 weight percent. Such a level of flame retardant resulted in an oxygen index of 32 to 34, 13 depending on thickness and density of the film, as can be seen above. The casting conditions and resulting properties 16 are summarized in Table II. പൂരുത്തിയും പ്രീത്യോ നിന്ന് (ജ്യാം വുടുക്ക് പോ പടിയുടുന്ന് ആ നന്നുന്ന വുടി വാഴി ആവസ്യം വ

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ఆమెక్కు గ్రామం పుర్మాణమైన కొండుకు కొండుకు ప్రముఖకారణ్ కొర్పుకు కొర్పుడు కొర్పుడు కొర్పుడు కొర్పుడుకు కొర్పుడు - ప్రారాహ్యం మూర్క్ కొండా కొర్పుడుకే ప్రముఖకాష్ట్రక కొరాయే. కోషం కొర్పుడుకోవడుకోవి మహీక కొర్వకోవారుకో ఉంది.

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ర్జుకార్యక్షామ్ కాండ్ కట్ట్ కొండాలు ఉందారు. ఎడ్డ్ కెక్కెస్ మీక్ కెక్కెస్ కెక్కెస్ ఈ మండ్రికి అయ్యాన్నారు. మాక్కెస్ కెక్ట్ స్టాన్స్ కెక్ట్ కెక్ట్ స్టాన్స్ కెక్ట్ స్టాన్స్ కెక్ట్ స్టాన్స్ కెక్ట్

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	Wet Film Thickness	18	19	
	Time In First. Drying Oven Minutes	13.0	10.4	వైద్య ప్రాంధికోండు. ఈ మైక ప్రాంత్రికోండి ప్రాంతించిన మండి ప్రాంతి ప్రాంతి ప్రాంతి ప్రాంతి ప్రాంతి ప్రాంతి ప్ర ప్రాంతి ప్రాంతి ప్రాంతి ప్రాంతి ప్రాంతి ప్రాంతి ప్రాంతి ప్రాంతి ప్రవాణ ప్రాంతి ప్రవాతి ప్రాంతి ప్రవాతి ప్రాంతి
	Oxygen Index	32.9	32.9	ිට ගිනිවේ ඉතිරිපතිවේ එමේ සම්බන්ධ සම්බන්ධ සිට එම සිට එම සිට එම සිට අතර අතු අතු මේ දැන්න එම සිට සිට එම සුලුපතිව අතු අතුම් අතු අතුම් සිට
. 5 27. . 3	Dielectric Strength volts/mil	1029,	965	ල ඉවස්වා ම විසින්ට ව ලබා සැබි විසින විසිට මිස්වා ට ටිස් වා ටිසිට දෙසුවුවෙනුව මැසිල්වස්ත පිට වෙදුසිස්විස් ට සිට ට ස්වාශ රාම්බයවෙමුණිකු සිසින්වු ලක්වෙස්ව වර්ණීට ට සිටිම් සිටිම්වෙස්ව සිටිම් සිටිම සිටි
TABLE II	Propagating Tear Strength Rm/mil	14.8	14.0	ముందు కింది కేంద్ర కేంద్ర కారా ఉంది కేంద్ర కేంద కార్పులు కేంద్ర కార్యాలు కార్యాలు కేంద్ర కారా కార్మాలు కేంద్ర కేంద్ర కేంద్ర కేంద్ర కేంద్ర కేంద్ర కేంద్ర కేంద్ర కేంద్ర కేంద్ర కేం
	Tensile Modulus psi	, 143,000 249,000	134,000	Talign Consider (1996) (1996) (1996) (1996)
	Dry Film Thickness mils	5.6	6.0	
2 12 .20 1 32.	Density Rm/cc		c 0.63 rol 1.3	ကြာလျှင်း ရေးက ပြုတ်လေးသည် သည်သို့ သို့သို့ရေးသည် သောသော လေးလိုင်းသည့် သို့သို့ရေးသည် သောသော လေးလိုင်းသည့် သော လေ့သည် သောလိုင်းသည် သည်သည် သည် သို့သည် သို့သည် သို့သည် သို့သည် သို့သည် လေ့သည် သည်သည် သို့သည် သည် သို့သို့သည် သည်သည် သည် သို့သည် သည်သည် သို့သည် သည်သည် သည်သည် သည်သည် သည်သည် သည်သည် သည
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Example 2

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A casting solution for an aromatic polyamide was 2 prepared by stirring a mixture of 16 grams of poly(imino-3 1,3-phenyleneiminocarbonyl-1,3-phenylenecarbonyl), 59 grams 4 of N,N-dimethyl acetamide and 4.4 grams of lithium chloride 5 which is employed so as to facilitate solubilizing the 6 polymer. A 25 mil wet film was cast onto a glass plate and 7 placed in a circulating air oven for 20 minutes and at a 8 temperature of 200°F so as to partially remove the solvent. 9 The clear film on the glass plate was then immersed in 10 water for one hour and thereafter dried at 350°F for three 11 hours. A tough, opaque, flexible, nonporous microcellular 12 film having good physical and electrical properties was 13 produced. 14 Example 3 15

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This Example is illustrative of method 1 described above wherein very low density, nonporous, microcellular film can be obtained by exposing the film to water vapor prior to any drying, i.e., the film is not partially dried before contact with water.

A 15% by weight solution of polysulfone (from ~21 Union Carbide), have the structure 22

was prepared using N,N-dimethyl formamide. The solution 26 was cast onto a glass plate and exposed to humid air until 27 a microcellular structure was formed as indicated by an 28 opaque off-white color. The film was removed after one 29 hour and dried at 150°C for one hour. The dried film was 30 nonporous, microcellular, flexible and tough. 31

Example 4 32

A solution of 10 grams of polyhydantoin (Bayer 33 4089) was dissolved in 40 grams of N,N-dimethyl formamide. 34

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A 20 mil thick wet film was cast onto a pyrex glass plate and put into an air swept drying oven and held at 250°F for five minutes. The partially dried clear film was immersed in water for four (4), hours. The now opaque film 5 was removed from the water and dried 30 minutes at 250°F, 30 minutes at 350°F and four hours at 400°F. The dried film was opaque, nonporous, microcellular, flexible and tough. THE STATE OF A THE STATE OF THE The microcellular structures of the invention can have incorporated therein a wide variety of small particle additives and/or fillers. The resulting structures 11 are relatively nonbrittle compared to a dense structure 12 containing a comparable amount of filler or additive. 13 Illustrative examples of additives are flame 14. retardants, antioxidants, pigments and the like. 15 The cellular structures will also be highly 16 useful as separators in fuel cells which do not utilize 17 alkaline electrolytes. The high temperature resistance, 18 strength and ability to be adhered to conductors such as 19 metals and carbon as well as the ease of electroplating a strong adherent metal film to the microcellular structures make them uniquely suitable for many fuel cell and battery component applications. Furthermore, the ability of the dense and cellu-24 lar materials described herein to both adhere tenaciously 25 to metals, carbon graphite, etc., substrates as well as 26 their high temperature solvent and corrosion resistance 27 to virtually all chemical substances except aprotic sol-28 vents and alkalies makes these polymers outstanding ma-29 terial for tank linings, pipe coatings and other thin film protective coatings. Their low permeability prop-31 erties (denser film) are also significant in this appli-. 32 33 cation. Another unique and highly useful application 34 of the microcellular film coatings, especially PPA, re-35 lies on the unusual low temperature performance of these 36 Thus, coatings for pipes and electrical cable 37 polymers.

1 conduit wraps can be used in extremely adverse low temperature, environments as low as about -268°C, with no adverse 3 effect. 22 This permits use with liquid helium and liquid mnitrogen without losing flexibility and with low dissipa-, tion factors. The mainter of the company was the This permits PPA (cellular film), to be used as insulating and protective materials in low temperature conductors. Such low temperature conductors are the clear 9 trend of the future and PPA should play an important role in these environments. the state of the s pactia las go tolum sait ಕಾಶ್ರಯ ಅಭ್ಯಾಪಕ್ಕ ಕಾರ್ಚಿಕ ಪ್ರಸ್ತಿಸಿಕ ಪ್ರಸ್ತಿಸಿಕ ಪ್ರಸ್ತಿಸಿಕ ಪ್ರಸ್ತಿಸಿಕ ಪ್ರಸ್ತಿಸಿಕ ಕಾರ್ಯಿಸಿಕ ಪ್ರಸ್ತಿಸಿಕ ಪ್ರಸ್ಟಿಸಿಕ ಪ್ರಸ್ತಿಸಿಕ ಪ್ರಸ್ತಿಸಿ ల సంముఖ్యమ్మాన్ని ఎక్కువాని కోమాన్ కోమాన్ని కార్యాలు కార్యాలు ముఖ్యమ్మాన్ మొఖ్యాలు ఉంది. మూర్తున్నా assistant the protection with the residual to the constitution రాగు ప్రత్యేఘ మేటూ కాడాన్ కొట్టు ఎక్కువ సమ్మాయ్లుకు కొట్టి మందుకుండాని. ార్మ్ ఉన్నాయి. కోమ్స్ కారార్ ఎక్కా గ్రామ్ చేశాల కోష ം അതുത്തിലെ ത്രണം വന്നത്തിലെ തുരുത്തിന്റെ വിവിധിക്കുന്നത്. വിവിധിക്കുന്നത് വിവിധിക്ക് വിവിധിക്കുന്നത്. ಕ್ಷಾ ಗ್ರಾಮಿಕ ಕರ್ನಾರಿಕಾಗಿಕರು ಆಗೆ ಬಿಂದು ಕೆಕ್ಟ್ ಎರೆ ಆ ್ಟ್ರಿಯಾ ಕಿತ್ರಕ್ಕೆ ಮಿನ್ನು ಬೆಳ್ಳಬಂದಿಕೆ ಅ ఆర్థ్యమ్మాన్ ఇండి ఈ నిజ్ఞంత్ర్మ్ ఈ అయిక్ ఎక్క్ స్ట్రీస్ చేశాకుండా చెప్పారు. ఈ ఆర్థ్యమ్మ్ ఈ ఆర్థ్యమ్మ్ ఈ ವಿಷ್ಠಾರ ಕ್ಷೇತ್ರವಾಗಿ ಕ್ರಾಮಾನ್ಯ ಕ್ರಾಮಾನ್ ಕ್ರಾಮಾನ್ಯ ಹೇಳಿದ್ದು ಪ್ರಾಮಾನ್ಯ ಕ್ಷೇತ್ರಗಳ ಕ್ರಾಮಾನ್ಯ Control of the State of the Control of the State of the S rolling on actions to the special terms of a communications. ្តីពីស្សារ ពេញសេសនា ពេញ ស៊ីដែល ២០ ១០ ខ្លាំ ១០ ២០១៣ ខេត្ត ១០០០០ ខែ ស្នាក់ ស្នាក់ ស្នាក់ ស្នាក់ ស្នាក់ ស្នាក់ ស្ में के पर्योग के में में किया है कि है है है है है है के का उन्हें की किया है है ಾರ್ಯವಾಗಿ ನಿರ್ದೇಶ್ವ ನಿರ್ವಹಣೆ **ಆಲ** ಅನ್ನು ನಿರ್ಮೇಶ್ವ ಗ್ರಾಮಕ್ಕೆ ನಿರ್ವಹಣೆ ಕಾರ್ಯವಾಗಿ ಕಾರ್ಯ and the second of the second o man nymetra ara-kan ingan-kanan dan basah dalah sakar dalah sakar dalah sakar dalah sakar dalah sakar dalah sa ు గృజ్ఞ గాహ్లామికియేకియోకాన్ని ఉద్దారం మేగా ఉన్నాయి. సమీప జ్ఞేగ్రామీకి మేగికి ఇంగాలా, ఎక్కువారి అన్న ఉంది. మూడికి మార్ట్ కెంక్ కెంక్ ఎక్కువ్స్ అన్నాయి. మూడికి మూడికి మూడికి మూడికి మూడికి మ రాగా కొడ్డా గ్రామం నాట్లు కొన్నారు. అదాకు సాండ్ కాటల్ కా ఉందినే ఉంది. కొన్నే

- 1. A shaped, relatively low density, microcellular solid polymeric article comprising a polymer
 selected from aromatic polyparabanic acids, aromatic polyhydantoins, aromatic polyamides, aromatic polyimides,
 aromatic polysulfones, aromatic polyimide-amides and
 combinations thereof, characterized in that the solid is
 made up of closed cells.
- 2. An article according to claim 1 characterized in that the article is a film.
- 3. An article according to claim 1 characterized in that the article is generally rod shaped with a relatively large length-to-diameter ratio.
- 4. An article according to claim 3 characterized in that said article is a fiber.
- 5. A thin coating of the article of claim 1 attached firmly to a substrate.
- 6. An article according to claims 1-5 characterized in that it has a density of about 0.3 to 1.5 g. per cubic centimeter.
- 7. An article according to claims 1-5 characterized in that it has a density of about 0.3 to 1.2 g. per cubic centimeter.
 - 8. A flexible circuit article characterized by:
 - (a) a structure according to claim 2; and
 - (b) a conductive circuit adhered to said film in a prearranged configuration.
- 9. A flexible circuit according to claim 8 characterized in that said conductive material is copper.
- 10. An article according to claim 8 or 9 characterized in that said conductive material has been electrodeposited on said film-shaped article.
- 2 11. An article according to claims 1, 2 and 5 or 6 characterized in that the surface thereof has been metal coated for decorative purposes.

- 12. An article according to claim 28 characterized in that said metal is chromium or aluminum.
- 13. An article according to claims 1-9 charac-1 terized in that the cellular surfaces have been coafed to with a thin layer of a catalytic material so that the resulting article is a supported catalyst structure. Same
- 14. An article according to claims 1-13 characterized in that said polymer is PPA.
- 15. An article according to claims 1-13 characterized in that said polymer is an aromatic polyhydantoin.
- 16. An article according to claims 1-13 charac-1 terized in that said polymer is an aromatic polyamide.
- . 17. A method of producing cellular structures according to claims 1-16 characterized by the steps of:
 - (a) forming a solution of said polymeric material in a solvent for said material;
 - casting said polymer solution onto a suitable surface to form an intermediate stage structure in any suitable prearranged configuration or extruding a fiber;
 - (c) exposing said resulting structure to an antibog of S.I orsolvent; by in message a seek at these his book
 - (d) precipitating solid polymer in the presence of said antisolvent;
 - (e) drying said precipitated solid; and
 - (f) recovering said shaped article.
 - 18. A method according to claim 17 characterized in that said antisolvent is water.
 - 19. A method according to claims 17 or 18 characterized in that said solvent is dimethylformamide.
 - 20: A method according to claims 17-19 characterized in that at least some solvent is evaporated from said intermediate stage structure prior to exposure of said structure to the antisolvent.

್ರಾಥ್ ಕ್ಷಾಗ್ರಾಕ್ಷ್ ನಿರ್ಣಾಪ್ತಿಯ ಕ್ರಾರ್ಡ್ ಕ್ರಾರ್ಡ್ ಪ್ರಾರ್ಥಿಸಿದ್ದಾರೆ.

- 21. A method according to claims 17-20 characterized in that said polymer is polyparabanic acid.
- 22. A method according to claims 17-21 characterized in that said suitable surface is a surface in which it is intended that the resulting structure will become permanently bonded.
- 23. A method according to claim 22 characterized in that said substrate is metallic.
- 24. A method according to claim 22 characterized in that said substrate is copper.
- 25. A method according to claim 22 characterized in that the substrate is glass.
- 26. A method according to claim 22 characterized in that the substrate is ceramic.
- 27. A method according to claim 22 characterized in that said substrate is a porous article.
- 28. A method according to claim 27 characterized in that said porous article is selected from wire, wood, paper, textiles, non-wovens and other plastics.
- 29. An article according to claims 1, 2, 6, 7, 14-16 is characterized in that it is paper-like in appearance and is coated with a photographic emulsion.
- 30. An article of clothing characterized by textiles prepared from fibers consisting of the article of claims 1, 3, 4, 6, 7, 14-16.
- 31. As a battery separator, a thin sheet of the article of claim 1, 2, 6, 7, or 14-16.
- 32. As a fuel cell separator, a thin sheet of the article of claim 1, 2, 6, 7, or 14-16.
- 33. A photographic paper according to claim 29 characterized in that said emulsion is a gelatin based emulsion.
- 34. A composite semi-conductor article characterized by a film made from a composition according to claims

1, 2, 6, 7, or 14-16 which has adhered to it a prearranged configuration of a metal oxide semi-conductor (MOS).

matrix reinforced with an article according to claims 1, 3, 6, 7, 14-16.

36. A shaped, relatively low density non-porous, microcellular polymeric article according to claims 1, 2, 6, 7, 14-16 characterized by being from about 1 mil to about 20 mils thick.

37. A shaped polymer according to claims 1, 2, 6, 7, 14-16 characterized in that it is from about 1 mil to about 20 mils thick and has a density of about 0.3 to about 1.5 g/cc, a tensile strength of about 5000 to about 10,000 psi, and an elongation of about 100% to 150%.

38. An article according to claim 37 further characterized by having a dielectric constant (10³ cps) of about 1.5 to 2.8 and dielectric strength of about 0.5 to about 3.0 KV/mil.

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ండునునుగా ఇద్ది అత్రికేస్తాడు. - జాగు **క్షాంక** అంగాలు అ**ద్దే మ**ర్యాలు కార్డ్ ఉ<mark>ది. - ఉద్దే</mark> - - - - ఇద్దం మార్క్ ఎండరమగు ఎండిమాక్కించగా ఎంది ఈ మంద్ర్ ఉద్దేశ్ మర్గులు ఉంది. మీడులో





EUROPEAN SEARCH REPORT

EP 80 30 1363

ategory	Citation of document with	ISIDERED TO BE RELEVANT indication, where appropriate, of relevant	Polessee	CLASSIFICATION OF TH APPLICATION (Int. Ct. 7)
	passages		Relevant to claim	
X	GB - A - 1 4	73 946 (EXXON)	1-38	C 08 J 9/28 5/18
A & 4. 1. 77	,	20. 20. 20.		G 03 C 1/76 G 11 B 5/70 B 01 D 13/04 C 08 J 5/04 H 01 L 21/31
			:	TECHNICAL FIELDS SEARCHED (Int. CL1)
·	·			C 08 J 9/28
	·	-	A C P	CATEGORY OF CITED DOCUMENTS : particularly relevant : technological background : non-written disclosure : intermediate document : theory or principle underlying the invention
			D	conflicting application document cited in the application citation for other reasons
	The present search re	port has been drawn up for all claims	8:	member of the same patent family,
of sea		Date of completion of the search 26-01-1981	Examiner	EMEESCH

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